Symptom burden, Metabolic profile, Ultrasound findings, Rhythm, neurohormonal activation, haemodynamics and health-related quality of life in patients with atrial Fibrillation (SMURF): a protocol for an observational study with a randomised interventional component

Emmanouil Charitakis,1 Ulla Walfridsson,1 Fredrik Nyström,2 Eva Nylander,3 Anna Strömberg,1 Urban Alehagen,1 Håkan Walfridsson1

ABSTRACT

Introduction: Atrial fibrillation (AF) is the most common cardiac arrhythmia, with an estimated prevalence of 1.5–2%. It is an independent risk factor for ischaemic stroke and is estimated to cause about 20–25% of all stroke cases. AF has a great impact on health-related quality of life (HRQoL); however, one unresolved issue related to AF is the wide variation in its symptoms.

Methods and analysis: The symptom burden, metabolic profile, ultrasound findings, rhythm, neurohormonal activation, haemodynamics and HRQoL in patients with AF (Symptom burden, Metabolic profile, Ultrasound findings, Rhythm, neurohormonal activation, haemodynamics and health-related quality of life in patients with atrial Fibrillation, SMURF) study is a prospective observational, cohort study, with a randomised interventional part. The aim of the study is to investigate, in patients with AF, the relationship between symptom burden and metabolic aspects, atrial function and different neurohormones, and the effect of radiofrequency ablation (RFA). The interventional part of the study will give an insight into the neurohormonal and intracardiac pressure changes directly after initiation of AF. Consecutive patients with symptomatic AF accepted for treatment with RFA for the first time at Linköping University Hospital are eligible for participation. The enrolment started in January 2012, and a total of 200 patients are to be included into the study, with 45 of them being enrolled into the interventional study with initiation of AF. The sample size of the interventional study is based on a small pilot study with 5 patients induced to AF while 2 served as controls. The results indicated that, in order to find a statistically significant difference, there was a need to include 28 patients; for safety reasons, 45 patients will be included.

Ethics and dissemination: The SMURF study is approved by the Regional Ethical Review Board at the Faculty of Health Sciences, Linköping, Sweden. The results will be presented through peer-review journals and conference presentation.

Trial registration number: NCT01553045;
Pre-results.

INTRODUCTION

Symptom burden, Metabolic profile, Ultrasound findings, Rhythm, neurohormonal activation, haemodynamics and health-related quality of life (HRQoL) in patients with atrial Fibrillation (SMURF) takes a broad view of symptoms and mechanisms of
atrial fibrillation (AF). In order to achieve this, we studied the quality of life, several neurohormones, metabolic aspects and atrial function in a randomised, controlled substudy where AF is induced: the ‘interventional SMURF-study’ was carried out.

AF is the most common cardiac arrhythmia with an estimated prevalence of 1.5–2% in the general population. However, a recent trial reports a considerably higher estimation of at least 2.9% in Swedish population. 1 2

AF is an independent risk factor for ischaemic stroke and is estimated to cause about 20–25% of all cases. 3 4 It is also associated with an increased risk of heart failure 5 and death. 6 Several clinical risk factors, ECG and echocardiographic features and biochemical makers are associated with an increased risk of AF. 7–9 Patients with AF with no or minimal heart disease and no other known non-cardiac causes are considered as ‘lone AF’ cases.

AF has a great impact on HRQoL. 10 11 One major unresolved issue in AF is the large variation in symptoms. In some patients, AF is diagnosed accidentally during routine check-up, while in others, symptoms are severe and disabling. There are data supporting the contention that change in AF episode duration, ventricular rate and AF cycle length can predict symptoms in patients with paroxysmal AF (PAF). 12 Moreover, there is evidence to show a relationship between perceived symptom burden and rhythm control, personality and gender. 13–15

Obesity, and thus insulin resistance, is a major contributor to cardiovascular (CV) disorder and hence adds to the incidence of AF. 16 17 Insulin resistance as a part of the metabolic syndrome is also linked to diastolic ventricular dysfunction 18 and might contribute to more pronounced symptoms of AF. 19 Thus, it is of interest to measure markers of insulin resistance such as obesity levels, fasting insulin, glucose and cholesterol in patients with AF.

Acute alcohol intake increases the risk of arrhythmias, but moderate intake of alcohol appears to harbour protective effects against CV and also other metabolic disorders. 21 High long-term intake of alcohol has been associated with an increased prevalence of AF, 22 but estimation of alcohol intake from questionnaires is affected by different sources of bias. Moreover, long-term intake of alcohol can be estimated by measuring ethyl glucuronide (EtG) in human hair. 23

Left atrial (LA) remodelling and enlargement is a powerful predictor of several CV events including stroke and death. 24 Using the area-length method from the apical 4 and 2 chamber views, it is possible to measure LA volumes (LAVs) in each phase of the atrial cycle and, furthermore, to quantify LA function by generating ejection fractions (EF) for each cycle. Two-dimensional speckle tracking (2D-SpTr) echocardiography is a relatively novel method for quantifying myocardial wall deformation using tracking of acoustic speckles. 25 It allows simultaneous analysis of radial and longitudinal myocardial deformation independent of insonation angle by automatically tracking myocardial segments and is validated against sonomicrometry. 26 This method may provide a comprehensive assessment of LA function. 27

AF is associated with the activation of several neurohormonal systems and, for some, the underlying activation mechanisms and their subsequent effects are unknown. The endocrine function of the heart is complex. B-type natriuretic peptide (BNP) and atrial natriuretic peptide (ANP) are examples of peptides involved in the cardiac endocrine. 28 Natriuretic peptides are mainly produced by atrial myocytes in normal participants. 29 As the heart is failing, an increased part of the production of ANP and BNP stems from the ventricular myocytes. Both peptides are known to be produced in response to the increased wall tension and in AF, in order to restore and maintain CV homoeostasis. 29–32

Elevated concentrations of these peptides have been reported in patients with AF without overt heart failure, 33 and these lead to natriuresis and vasodilation in order to normalise haemodynamics. However, it has been reported that prolonged AF is associated with a reduced capacity by the atria to produce natriuretic peptides. 34 The plasma concentrations of the natriuretic peptides are elevated in patients with AF and are known to decrease after conversion to sinus rhythm (SR). 35 BNP and its N-terminal fragment of proBNP are not only used in risk assessment of patients with heart failure but also in those with AF. 36 Owing to its instability, ANP has not been used in clinical routine. A new, more stable method for quantification of the mid-regional fragment of the N-terminal precursor of proANP (MR-proANP) has been introduced, 37 and may be useful in risk stratification of patients with AF. Vasopressin (AVP), a non-cardiac plasma marker of CV disease, is released from the neurohypophysis in response to changes in plasma osmolality, and is involved in osmoregulation and CV homoeostasis. The plasma concentration of AVP is increased in patients with heart failure and is associated with left ventricular dysfunction. 38 However, AVP, as such, is not a clinically useful plasma marker because it rapidly degrades in the circulation. Instead, copeptin, the C-terminal fragment of proAVP, has emerged as a promising surrogate biomarker for the measurement of AVP concentration. 39 Adrenomedullin (ADM) originates primarily in adrenal medulla and endothelial cells where cellular stress, ischaemia and hypoxia induce increased production. 40 It is capable of promoting vasorelaxation, natriuresis and diuresis, thus contributing to maintaining circulatory homoeostasis. 41 The mid-regional portion of pro-ADM (MR-proADM) is more stable than ADM and therefore better suited for clinical practice and assessment in stored samples. 42 The reaction to AF initiation is unknown. Moreover, the role of ADM and AVP-hormonal system has not been studied in patients with AF.
AIMS
The aims of this study are: (1) to correlate the variety in symptom burden in patients with AF eligible for radiofrequency ablation (RFA) as measured by patient-reported outcomes: HRQoL, perceived control over the heart disease, anxiety and depression with objective indicators such as peptide levels, echocardiographic data, haemodynamics and metabolic profile; (2) to study the short-term effects of successful RFA peptides, LA function and patient symptoms; (3) to study the level of different peptides in peripheral vein blood and two sites of the heart (coronary sinus (CS) and LA) and to give an insight into the neurohormonal reaction after RFA in patients with AF; (4) to echocardiographically quantify LA function and intracardiac pressures in patients with AF; and (5) to analyse relationships of components of the metabolic syndrome with increased risk for recurrence of AF and increased LA V.

Furthermore, in order to study possible neurohormonal and intracardiac pressure changes directly after initiation of AF, an interventional study is to be performed.

METHODS
Study design
This is an observational study in a single-centre cohort. A group of patients will also be included in an interventional randomised substudy. The first patient was included in January 2012. A total of 200 patients are planned to be included into the main study and 45 patients into the interventional study with a follow-up period of 12 months. No participants will be recruited into the randomised section of the study until prior to registration at http://www.clinicaltrials.gov. Enrolment is expected to last approximately 2.5 years and the study is to be completed by the end of 2015.

Study population
Patients eligible for study participation are those referred for RFA of AF at the University Hospital in Linköping, Sweden. Consecutive patients are asked about study participation and are included if they meet the inclusion and do not have any exclusion criteria. Inclusion and exclusion criteria are presented in table 1.

Patients eligible for the main study, who are free from arrhythmia during 4 days prior to RFA, can be included in the interventional study. The recruitment process and follow-up is presented in figure 1.

Informed consent and ethical considerations
All patients gave written informed consent, and the study participation is documented in the patient’s medical record. Informed consent will be obtained by medical doctors participating in the study. The study complies with the Declaration of Helsinki.43 The study is to be periodically externally monitored in order to identify potential sources of bias. Any changes of study objectives, protocol and population, will require a formal amendment to the protocol. All study-related information is stored securely in a local database. All laboratory specimens and other data will be marked by a coded identification number. Further use of stored blood samples will require renewed evaluation by the Regional Ethical Review Board at the Faculty of Health Sciences, Linköping, Sweden. All recorded personal identifiers will be stored in the investigator site file and identified by code number. All investigators will be given access to a cleaned data set. All data sets will be password protected. To ensure confidentiality, data dispersed to project team members will be blinded of any identifying participant information. The results of the trial will be presented at regional and international congresses, and in peer-reviewed journals, and will also be released to the participating physicians, referring physicians and patients.

Questionnaires
Patient-reported outcome measures (PROMs) are assessed with four questionnaires, at the start of the study (baseline measurement) and after 4 and 12 months. The following questionnaires are used.

Medical outcomes study 36-item short-form health survey
36-item short-form health survey (SF-36) is a generic questionnaire designed to measure an individual’s physical and mental health (MH). It comprises 35 items grouped into eight scales and one question concerning changes in health outside the scales. The eight scales represent physical functioning, role limitations due to physical health problems, bodily pain, general health, vitality (energy/fatigue), social functioning, role limitations due to emotional problems (RE) and MH (psychological distress and psychological well-being). The eight scales are summarised in two dimensions; physical and

<table>
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<th>Table 1</th>
<th>Inclusion and exclusion criteria in the SMURF main study</th>
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<td>Inclusion criteria</td>
<td>Exclusion criteria</td>
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<td>Patients &gt;17 years of age with paroxysmal or persistent AF, referred to the Linköping University Hospital for the first RFA for AF</td>
<td>Previous catheter or surgical AF ablation</td>
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<td>Informed consent</td>
<td>Ejection fraction &lt;35%</td>
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AF, atrial fibrillation; RFA, radiofrequency ablation.
mental component summary. For each of the eight scales and the two summary dimensions, scores are coded, summed and transformed to a scale from 0 (worst possible health) to 100 (best possible health). The scoring of the SF-36 data is carried out as described by Ware and Sherbourne, and Ware et al. SF-36 has been widely used in research including in studies on patients with arrhythmias.

Arrhythmia-specific questionnaire in tachycardia and arrhythmia

The disease-specific arrhythmia-specific questionnaire in tachycardia and arrhythmia (ASTA) questionnaire is divided into three separate parts, where ASTA part I evaluates the patient’s latest episode of arrhythmia, pharmacological treatment and—important at follow-up—one question concerning the existence of any arrhythmia. ASTA part II covers symptom burden: the ASTA symptom scale has nine items and a four-point response scale with response alternatives from 0 to 3: ‘No (0); Yes, to a certain extent (1); Yes, quite a lot (2) or Yes, a lot (3)’. A summarised score is calculated for the items, a higher score implies higher symptom burden due to the heart rhythm disturbance. Outside of the symptom scale there are questions concerning frequency of arrhythmia, duration and palpitations, and if there are factors influencing arrhythmia occurrence and the experience of near syncope and/or syncope in connection with

Figure 1 Enrolment and follow-up flow chart (AF, atrial fibrillation; HRQoL, health-related quality of life; RFA, radiofrequency ablation).
arrhythmia with the response alternatives ‘No or Yes’. ASTA part III is on HRQoL: the ASTA HRQoL scale has 13 items and describes the influence of arrhythmia on daily life. The response alternatives are the same as for the ASTA symptom scale (please see above) where higher scores reflect a worse effect on HRQoL.47 48

Hospital anxiety and depression scale
The domain-specific hospital anxiety and depression scale (HADS) questionnaire is used to evaluate symptoms of anxiety and depression. The questionnaire consists of two sections, where anxiety is assessed with seven questions (HADS-A) and depression with the remaining seven (HADS-D). Responses are scored on a scale of 0–3; with higher scores denoting more psychological distress. The score for each subscale (anxiety and depression) can range from 0 to lowest as a maximum score of 21. The scores are categorised as normal (0–7), mild (8–10), moderate (11–14) and severe (15–21) anxiety and/or depression, respectively.49 50

Control attitudes scale
The control attitudes scale (CAS) is developed to measure perceived control among patients with cardiac diseases. In this paper, the validated Swedish version is used. The scale consists of four items, two about perceived control and two about helplessness. Furthermore, two of the items reflect the patient’s own perceptions and two reflect the patients’ perception about family members’ perceived control. Patients rate each item on a seven-point Likert-type scale, from ‘Not at all’ to ‘Very much’. Two of the item scores are reversed and then a total score is calculated by summing the responses. The total score can vary between 4 and 28, where higher scores indicate greater perceived control.51 52

Interviews
A subset of patients will be included in an interview study exploring the patients’ daily life situation due to AF. An interview-guide including semi-structured questions is constructed. All interviews are conducted by one researcher and begin with an initial open-ended question followed by questions intended to guide, refocus or clarify the respondent’s answer. The interviews will be made before RFA, and will be audio recorded with the patients’ permission. Qualitative inductive content analysis (conventional content analysis) is chosen to interpret the meaning from the content of text data by classifying the data into codes and categories. The data will be coded aided by the software system NVivo9 (QSR International Pty Ltd, Melbourne, Australia).

Brief intermittent transtelephonic ECG
The ECG system used has been developed by Zenicor AB and consists of a device that can register a short ECG sequence (30 s) by having participants place their thumbs on two measuring plates. The method has been validated53 and used in order to improve screening of silent AF in patients with ischaemic stroke54 and as an event-recorder.

A registration is sent by telephone to a central database, which can only be accessed by authorised personnel. Rhythm analysis is performed prior to randomisation on patients who are eligible for the interventional study. The rhythm is classified into the following categories: SR, AF, atrial tachycardia or ‘not specified rhythm’.

Echocardiography
Before RFA, all patients undergo a transthoracic and a transesophageal echocardiographic examination (TTE and TEE, respectively) using the GE Vivid 7 or Vivid E9 system (GE Healthcare, Horten, Norway) with a 3.5 MHz transducer for TTE and a 7 MHz transducer for TEE. Measurements and evaluations are performed according to the guidelines of European Society of Echocardiography.55 Blood pressure is recorded and heart rate monitored during the examination. All data are stored digitally for further offline analysis (Echo PAC version BT 12, GE Healthcare, Horten, Norway).

TTE measurements
Left ventricular EF is calculated by biplane Simpson’s method. LAV is measured using the biplane area-length method and is corrected for body surface area to obtain LA index (LAVI). Left ventricular filling velocities (E and A) are obtained by pulsed-wave Doppler in the apical four-chamber (4CH) view. The ratio of E/A velocity is measured. Tissue Doppler images in the 4CH view are obtained and used offline in order to measure the early diastolic velocity (E’) in the mitral septal annulus. E/E’ ratio is then calculated.

Transesophageal echocardiography
Patients undergo TEE examination under pharyngeal anaesthesia. The LA appendage (LAA) is visualised and examined thoroughly for the possible presence of thrombus. Filling and emptying LAA flow velocities are assessed by pulsed wave Doppler. The sample area is positioned at the outlet of LAA, 1 cm away from the LA cavity.

Speckle tracking echocardiography
2D echocardiographic images of the LA are obtained from the apical 4CH and 2CH view for speckle tracking evaluation. All images are obtained with a stable ECG recording. Three consecutive beats are stored in a cine loop format. The frame rate is adjusted between 55 and 100 frames/s. The endocardial border of the LA is manually traced. A region of interest (ROI) is automatically generated by the software (Echo PAC version BT 12, GE Healthcare, Horten, Norway); manual adjustments are performed if necessary. Subsequently, the ROI is divided into six segments and if these are of acceptable quality then the software is allowed by the user to perform further analysis. Segments of poor quality are either automatically or manually rejected from further processing. Finally, strain and strain rate curves are
generated. Calculations by averaging the values observed in all LA segments are made for the measurement of peak atrial longitudinal strain, peak atrial contraction strain, time to peak longitudinal strain, time to peak contraction strain, LA strain rate during ventricular systole (LA-SRs), strain rate during ventricular passive filling (LA-SRE) and strain rate during atrial active contraction (LA-SRA). If some segments are excluded from the analysis, the calculation is to be made by averaging the remaining segments.

Ablation procedure

Procedures are performed under conscious sedation using propofol and remifentanil. Two trans-septal sheaths are inserted through the right femoral vein into the LA and perfused using heparinised saline with irrigation rate of 3 mL/h. Heparin is administered to maintain an activated clotting time of >350 s throughout the procedure. RFA is performed under the guidance of a computer-based mapping system, CARTO ( Biosense Webster, Diamond Bar, California, USA). Mapping and ablation are performed using an open-irrigated catheter (ThermoCool, Biosense Webster, Diamond Bar, California, USA). A 7-F, 20-pole circumferential diagnostic catheter is used for the assessment of pulmonary vein activation and isolation (Lasso, Biosense Webster, Diamond Bar, California, USA). Radiofrequency energy is delivered in a power-controlled mode with maximum energy setting of 35 W at an irrigation rate of 17–30 mL/min, and the maximum energy setting in the posterior wall is 25 W. The end point of the procedure is electrical disconnection of all pulmonary veins by antral ablation and is defined during SR by entry and exit block of all pulmonary veins, that is, DC conversion is performed if needed prior to the end of the procedure. In patients with persistent AF, additional ablation in order to create LA lines is at the discretion of the operator and verified by pacing manoeuvres.

Pressure measurements

The sagittal thoracic diameter is measured in the second intercostal space and the reference pressure (zero level) is placed in the middle of this diameter. A-wave, v-wave and mean pressures are measured in the right atrium (RA), LA and systolic and diastolic pressures in the right ventricle (RV) using the multipurpose (MR A1) high flow (Cordis) 5F catheter, during quiet breathing, and recorded at least for 15 s for offline analysis (EP-WorkMate; St Jude Medical, Saint Paul, Minnesota, USA).

Markers of metabolic syndrome and measurement of long-term alcohol intake

The laboratory analyses are performed at the Department of Clinical Chemistry at Linköping University Hospital. Levels of cholesterol, high-density lipoprotein (HDL) and triglycerides are measured using enzymatic methodology and spectrophotometry (Selectra E, Vital Scientific, the Netherlands/Triolab). Low-density lipoprotein (LDL) is calculated by Friedewalds formula: LDL=cholesterol−HDL−0.45×fS/P-triglycerides. Glycated haemoglobin (HbA1c) is analysed according to the Swedish Mono-S high performance liquid chromatography (HPLC) method, and all HbA1c values are converted to diabetes control and complications trial (DCCT) standard values using the formula: HbA1c DCCT=0.923×HbA1c (Mono-S)+1.345 (R²=0.998) and are to be reported in international federation of clinical chemistry and laboratory medicine (IFCC) units. Levels of apolipoprotein B (apoB) and apolipoprotein A1 (apoA1) are measured by immunoturbidimetric assays (Advia 1800, DakoCytomation, Glostrup, Denmark).

Hair samples are collected as close as possible above the skin and the proximal 2 cm are used for estimation of amount of EtG. Analyses are performed as published earlier.25

Cardiac biomarkers

Blood samples are collected in plastic vials containing EDTA. The vials are centrifuged at 3100 g for 20 min and then frozen at −70°C. No sample is to be thawed more than twice.

Concentrations of proBNP 1-76 (NT-proBNP) are measured on the Elecsys 2010 platform (Roche Diagnostics, Mannheim, Germany). Plasma concentrations of MR-proADM and copeptin are measured on a kryptor platform (Brahms AG, Hennigsdorf, Germany) and MR-proANP in commercial assay (MR-proANP; Brahm AG, Hennigsdorf, Germany).

Study’s flow chart

First contact

After screening, potentially eligible patients receive written information about the study and the following forms: SF-36, ASTA, HADS and CAS. Patients eligible for the interventional study are provided with a transtelephonic device and instructed to record and send ECG recordings twice daily and extra recordings in case of symptoms during the 4 days prior to RFA.

Baseline evaluation

Prior to RFA patients receive further oral information and informed consent is signed. A full baseline evaluation, including medical history, physical examination and a 12-lead ECG, is performed. All patients undergo TTE, TEE and a CT scan of the heart according to the clinical routine.

All registered transtelephonic ECGs sent from patients eligible for the interventional study are screened, and if free from arrhythmias, the respective patient can be included, otherwise the patient continues in the main study (figure 2).

The sagittal abdominal diameter is measured and a tuft of hair is cut for the analysis of EtG.
All patients are then catheterised according to clinical routine and blood samples (glucose, HbA1c, insulin, blood status, aspartate aminotransferase, alanine aminotransferase, lactate dehydrogenase, bilirubin, cholesterol levels including LDL, HDL, lipoproteins A and B, creatinine, cystatin C, thyroid-stimulating hormone, T4, T3 and high-sensitivity C reactive protein) are obtained from the femoral vein. Blood samples for the analysis of NT-proBNP, MR-proANP, copeptin and MR-proADM are drawn from the femoral vein, CS and LA (baseline sampling) using a multipurpose (MR A1) high-flow (Cordis) catheter. Intracardiac pressures are recorded in the RA, LA and RV.

Interventional study: induced AF
After baseline blood sample retrieval and intracardiac pressure measurements, patients randomised to the active group are induced to AF with burst pacing from CS (CL: 170–300 ms). AF is maintained for 30 min and immediately re-induced when necessary. Blood samples for the analysis of NT-proBNP, MR-proANP, copeptin and MR-proADM are drawn from the femoral vein, CS and LA (baseline sampling) using a multipurpose (MR A1) high-flow (Cordis) catheter. Intracardiac pressures are recorded in the RA, LA and RV.

Follow-up
The day after RFA, blood samples are drawn from a peripheral vein for peptide analyses. At 4-month follow-up, 24 h Holter, blood sampling and assessment with the previously used questionnaires are performed. TTE control is performed in a subgroup of patients who are residents of the Östergötland County. At 12-month follow-up, only assessment with questionnaires will be made.

End points
Primary end point
Primary end point is HRQoL and arrhythmia symptom burden in relation to the level of four different peptides: NT-proBNP, copeptin, MR-proADM and MR-proANP.

Interventional study, primary end point
Primary end point for the interventional study is changes in NT-proBNP, MR-proANP, MR-proADM and copeptin 30 min after the induction of AF compared to controls in three sample sites (femoral vein, LA and CS).

Secondary end points
- Reaction of NT-proBNP, MR-proANP, copeptin and MR-proADM directly after 18 h and 4 months after RFA in relation to heart rhythm at admission, heart rate and type of AF.
- Measurements of a possible concentration of copeptin and MR-proANP indicating a possible cardiac production of the two biomarkers and the levels of NT-proBNP and MR-proANP in LA and in CS.
- Correlation between MR-proANP, NT-proBNP, copeptin and MR-proADM levels, and LA function assessed by 2D-strain measurements.
- Echocardiographic description of LA function by the assessment of LAV, LA EF, 2D strain, and strain rate of the LA and LA appendix function in relation to heart rhythm, type of AF and long-term alcohol consumption.
- Correlation between estimated alcohol consumption, HRQoL and arrhythmia freedom after RFA.
- Correlation between the metabolic syndrome, sagittal abdomen diameter and AF recurrence after RFA as well as to LAV.
- HRQoL and symptom burden in patients with AF prior to, and 4 and 12 months after RFA.
- Peptide levels and LA function as predictors of arrhythmia-freedom 4 months after ablation.
- Effect of RFA on LA, LAA function and peptide levels 4 months after the ablation.
- Levels of NT-proBNP, MR-proANP, MR-proADM and copeptin in peripheral blood, LA and CS in relation to different types of AF and to CHA2DS2-VASc, and AF risk score.
- RA, RV and LA pressures in patients with AF eligible for RFA in relation to LA function and patients’ symptom burden.
- Pressure changes in RA, RV and LA after induction of AF.
- A description of how a patient’s daily life is influenced; exploring the relationship between arrhythmia-specific symptoms, HRQoL, anxiety and
depression, perceived control and healthcare contacts, a qualitative study.

Interventional study, secondary end points

- Activation of neurohormonal systems and changes in intracardiac pressures after initiation of AF in patients with a high symptom burden.
- Correlation between peptide changes, and changes in recorded pressures and echocardiographic data.

Randomisation concerning the interventional study

Forty-five patients are to be randomised. Randomisation is performed with a 2:1 allocation ratio, that is, for every two patients who are randomised to AF induction, one is to be randomised to the control group. Carefully prepared, sealed, opaque, identical and serially numbered envelopes produced by Linköping’s academic research centre are provided in order to ensure impartial randomisation. A trial investigator will assign participants to interventions. Owing to the nature of the interventional study, blinding of participants or staff was not possible.

Statistical issues and sample size consideration

Baseline characteristics will be reported as means and SD for parametric and median, and IQR for non-parametric data. Normality will be tested by Kolmogorov-Smirnov test with p<0.05 to be considered as statistically significant. Continuous data that are not normally distributed will be transformed if needed. Categorical data will be presented as number and percentage. All tests performed will be two-sided and a p<0.05 will be considered statistically significant.

Full statistical analysis is to be published separately when presenting each outcome. STROBE statement for the main study and CONSORT statement for the interventional study will be followed when publishing the results.

All missing values and those lost to follow-up will be excluded from the respective analysis. The analyses will be performed using SPSS V.22.0 (SPSS, Chicago, Illinois, USA). To the best of our knowledge, there have been no previous studies performed addressing our primary and several of the secondary end points, thus a power analysis was not possible. There have been studies on the reaction of ANP and BNP in patients with AF after RFA, and after successful electrical conversion to SR in which 40–60 patients were included, in order to show significant results. We concluded that 200 patients would be sufficient. The interventional study was preceded by a small pilot study where five patients were induced into AF and two patients served as controls. The assumption of the possible difference on MR-proANP level was that as MR-proANP is prereproduced and stored in cardiac myocyte granules, the initiation of AF could lead to a direct release of the peptide and higher levels of MR-proANP could be measured 30 min after the induction to AF. The results indicated that, in order to find a statistically significant difference between the intervention and the control group on the reaction of the levels of MR-proANP with a power of 90% and type I error of 0.05, there was a need to include 28 patients. As there are other neurohormonal systems to be tested, we increased the number of participants to 45 for safety reasons. The sample size calculation was performed using a commercially available statistical analysis software package (STATISTICA 10; Statsoft, Dell Statistica, Tulsa, Oklahoma, USA). Unequal allocation ratio was taken into account.

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Contributors

EC participated in all study aspects, design, pre-study administration, performing echocardiography and in laboratory analyses, drafted, and revised and approved this manuscript. UW participated in the study design, created the ASTA questionnaire for analysis of symptom burden and health-related quality of life, was responsible for analysing data on symptom burden and health-related quality of life, was responsible for the associated part of the manuscript, and revised and approved this manuscript. FN participated in the study design with special responsibility for data on the metabolic syndrome, and revised and approved this manuscript. EN participated in the study design with special responsibility for data on echocardiography and intracardiac pressures, and revised and approved this manuscript. AP participated in the study design, was co-responsible, with UW, for analysis of symptom burden and health-related quality of life, and revised and approved this manuscript. UA participated in the study design with special responsibility for data on neurohormones, and revised and approved this manuscript. HW conceived the idea of the study, is the main investigator with overall responsibility for the study, and revised and finalised this manuscript.

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Competing interests

None declared.

Ethics approval

The Regional Ethical Review Board at the Faculty of Health Sciences, Linköping, Sweden.

Provenance and peer review

Not commissioned; externally peer reviewed.

Data sharing statement

The results will be presented through peer-review journals and conference presentation. Additional publications are planned on this data from this institution.

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